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(S) Sulphonated polyarylethersulphone copolymers and process for the manufacture thereof.

(57) A polyarylethersulphone copolymer which is a hydrophilic sulphonated copolymer derived by controllably sulphonating a copolymer having the repeat unit of formula

together with the repeat unit B of formula

substantially all the units A being sulphonated after sulphonation and substantially all the units B remaining nonsulphonated after sulphonation. The sulphonation is effected

using 98% w/w sulphuric acid.

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SULPHONATED POLYARYLETHERSULPHONE COPOLYMERS

The present invention relates to certain sulphonated polyarylethersulphone copolymers.

We have found that polyarylethersulphone polymers containing only the repeat units of formula

dissolve in concentrated sulphuric acid (98% w/w) and sulphonate very rapidly to give completely water-soluble products, probably due to the substitution of -SO<sub>2</sub>OH groups in all or most of the aromatic rings in the subunits

of the polymer (see Examples 7 to 9), monosulphonation occurring in the case of the polymers with ortho or para disposed ether linkages in the sub-units and disulphonation occurring in the polymers with meta disposed linkages in the sub-units.

By contrast we have found that the polyarylethersulphones containing only the repeat unit of formula

are virtually non-susceptible to sulphonation in concentrated sulphuric acid (98% w/w), the sulphuric acid dissolving the polymer but not changing it, (see Example 10), even though other sulphonating agents such as oleum and chlorosulphonic acid, and even a mixture of concentrated sulphuric acid with a little oleum, completely sulphonate the polymer and/or degrade it (see Examples 11 and 12). Consequently polyarylethersulphone

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copolymers containing repeat units A and B may be controllably sulphonated in concentrated sulphuric acid (98% w/w) to give hydrophilic sulphonated copolymers (ranging up to completely water soluble polymers) by varying the proportion of the repeat unit B in the copolymer.

According to the present invention there is provided a polyarylethersulphone copolymer which is a hydrophilic sulphonated copolymer derived by controllably sulphonating a copolymer having the repeat unit of formula

together with repeat unit of formula

substantially all the units A being sulphonated after sulphonation and substantially all the units B remaining non-sulphonated after sulphonation.

The polyarylethersulphone copolymer of the invention is preferably one derived by sulphonating a copolymer which has 1 to 99 mole % of units A and 99 to 1 mole % of units B, and in particular a copolymer which has 5 to 80 mole % of units A and 95 to 20 mole % of units B.

The control of the sulphonation reaction should be such that the resulting copolymer preferably has a degree of hydrophilicity corresponding to a water absorption capacity at ambient temperature of about 2 weight % water absorption to complete solubility in water.

The hydrophilic polymers of the invention (which preferably absorb 2-40 weight % water, particularly 5-30 weight % water, at ambient temperature) are potentially useful as membrane materials, e.g. for ultra-filtration

processes, e.g. desalination and removal of microorganisms, since they are not only hydrophilic in character but retain considerable strength even when containing a significant quantity of water (e.g. up to 20 5 weight \* water) (see Example 5).

Ionomers of the above-defined copolymers which are sulphonated with -SO<sub>2</sub>OH groups may be readily prepared e.g. by converting the -SO2OH groups to salts such as  $SO_3-M^+$  (where M = an alkali metal or NR<sub>4</sub>, R being an 10 alkyl group); these too have utility as hydrophilic polymers. Accordingly, the sulphonyl groups of the copolymers of the invention preferably have the formula -SO<sub>2</sub>Y where Y is H, an alkali metal, or NR<sub>4</sub> where R is an alkyl group.

Copolymers having the repeat units A and B may be conveniently prepared by condensation of the appropriate dihydroxy phenol (e.g. hydroquinone, catechol or resorcinol), 4,4'-dihydroxydiphenylsulphone and 4,4'dichlorodiphenylsulphone, and an alkali metal carbonate or 20 bicarbonate in the presence of a sulphone or sulphoxide solvent, using the method of preparing polyarylene polyethers described in Canadian Patent 847 963.

The present invention is now illustrated by the following examples.

EXAMPLE 1 25

A copolymer (20 gms) of repeat units A (para disposed ether linkages in sub-units) and B containing 25 mole % of units A (and correspondingly 75 mole % of units B) was shaken for 18 hours with concentrated sulphuric acid (98% 30 w/w) (110 ml) to give a very viscous solution. Further (50 ml) concentrated sulphuric acid was added and shaking continued for a further 18 hours (i.e. 36 hours in all).

This solution was then poured into distilled water (800 ml) in a Waring blender, to give a white precipitate 35 which was filtered and washed 3 times with fresh water in

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the blender, being separated by filtration after each wash.

The product was dried overnight at about 65°C in a vacuum oven.

Copolymers of repeat units A (para disposed ether 5 linkages in sub-units) and C of different composition (varying from 25 mole % A to 66.7 mole % A and correspondingly 75 mole % B to 33.3 mole % B) were used in this Example. All the prepared products after analysis by 10 220 MHz nmr spectroscopy indicated monosulphonation on substantially all of the sub-repeat units

in the polymer chain but no sulphonation on any of the repeat units B. All the products were soluble in dimethyl 15 formamide and dimethylsulphoxide.

#### EXAMPLE 2

A copolymer (5 g) of repeat units A (para disposed ether linkages in sub-units) and B containing 80 mole % A and 20 mole % B was shaken overnight with concentrated 20 sulphuric acid (98% w/w) (20 ml). Further (20 ml) concentrated sulphuric acid was added and shaking continued for a further 24 hours. This solution was then poured into distilled water in a Waring blender to give a white precipitate which was filtered and washed 4 times with water, and dried overnight in a vacuum oven.

Copolymers of repeat units A (para disposed ether linkages in sub-units) and B containing respectively 5 mole % A/95 mole % B, 90 mole % A/10 mole % B, 40 mole % A/60 mole % B were sulphonated in a similar fashion to the copolymer 80 mole % A/20 mole % B, although starting from different quantities of copolymer and using different amounts of concentrated sulphuric acid.

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The products were analysed by 220 MHz spectroscopy and indicated monosulphonation on substantially all of the sub-repeat units

5 in the polymer chain but no sulphonation on any of the repeat units B.

# EXAMPLE 3

The sodium salts of the sulphonated copolymers of Example 2 were prepared by neutralisation in excess NaOH solution. The neutralisation of the sulphonated 10 mole % A/90 mole % B copolymer is described as a typical example.

The sulphonated 10 mole % A/90 mole % B copolymer of Example 2 (50 g) was stirred overnight in a solution of NaOH (30 g) in water (600 ml). The mixture was then heated at 60-80°C all day and cooled to 40°C. The product was filtered, washed 4 times with water and oven dried.

The sodium contents of the sulphonated 10 mole % A/
20 mole % B, 20 mole % A/80 mole % B and 40 mole % A/
60 mole % B copolymers were determined using flame
emission spectroscopy (FES) and confirmed the nmr
spectroscopic analysis indicating that substantially all
the units A had been monosulphonated. The results were as
follows:

25	Molar Ratio A/B in copolymer	Na content of sulphonated copolymer found by FES	Monosulphonation of units A requires
	10/90	0.53%	0.58%
	20/80	1.20%	1.00%
	40/60	2.30%	2.06%

# EXAMPLE 4

The water absorption of the copolymers of Example 2 before sulphonation and after sulphonation, and also of the neutralised sulphonated copolymers of Example 3, were 5 measured after steeping initially dry films of the copolymers (cast from solution in dimethyl formamide) in water for 24 hours at ambient temperature. The results were as follows:

Molar Ratio A/B in copolymer	Water absorption (weight %) of copolymer before sulphonation (Example 2)	Water absorption (weight %) of sulphonated copolymer (Example 2)	Water absorption of (weight %) of neutralised sulphonated copolymer (Example 3)
5/95	2.4	4.5	2.4
10/90	2.5	4.4	3.6
20/80	2.3	4.5	5.5
40/60	3.9	8.9	13.1
80/20	not measured	not measured	36.6

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#### EXAMPLE 5

The sulphonated 40 mole % A/60 mole % B and 80 mole % A/20 mole % B copolymers of Example 2 were tested in the form of cast film for mechanical properties, both dry and after absorption of moisture. For comparison purposes a homopolymer containing units B only (not sulphonated) was also tested. The results were as follows:

Polymer Used		Water Content (wt %)	1% Secant Modulus (NMm <sup>2</sup> )	Yield Stress (NMm <sup>2</sup> )	Tensile Strength (NMm <sup>2</sup> )	Elon- gation at break (%)
Homo- polymer of	Dry	<del>-</del>	1450	<b>-</b> .	74 ·	8.2
units B	Wet	1.0	1350	-	-	-
Sulphon- ated copolymer 40 mole % A/ 60 mole % B	Dry Wet		1700 1500	68 45	62 -	10.1
Sulphon- ated copolymer 80 mole % A/	Dry		1750	59 32	<b>4</b> 7 26	14.7 32.1
20 mole % B	Wet	19.5	1030	32	20	J4 • I
( - indica	ites	not measu	ured)			

### EXAMPLE 6

A copolymer (5 g) of repeat units A (meta disposed ether linkages in sub-units) and B containing 40 mole % A and 60 mole % B was dissolved in concentrated sulphuric acid (98% w/w) and left for 36 hours. This solution was poured into distilled water to give a precipitate which was filtered, washed 3 times with water and oven dried. The product was analysed by 220 MHz spectroscopy and indicated disulphonation on substantially all of the sub-repeat units

in the polymer chain but no sulphonation on any of the repeat units B.

The water absorption of the copolymer was measured after steeping an initially dry film of the copolymer (cast from solution in dimethyl formamide) in water for 24 hours at ambient temperature; the value obtained was 7.5 weight %.

#### EXAMPLE 7

A homopolymer of repeat unit A (para disposed ether linkages in sub-units) (10 g) was dissolved in concentrated sulphuric acid (98% w/w) with stirring and samples (50 ml) removed from time to time and precipitated in dilute sulphuric acid or saturated sodium sulphate; the polymer samples corresponding to the various times of sulphonation were analysed by 220 MHz nmr spectroscopy as solutions in dimethyl sulphoxide and found to have the following properties:

1 hour .85% of the units bore an -SO<sub>2</sub>OH group (isolated as Na salt).

2 hours 99% of the units bore an -SO<sub>2</sub>OH group (isolated as Na salt).

20 4.5 hours 100% of the units bore an -SO<sub>2</sub>OH group (isolated as Na salt).

23 hours 98% of the units bore an -SO<sub>2</sub>OH group (isolated as free acid).

Substantially all the sub-units bore one -SO2OH group.

25 EXAMPLE 8

A homopolymer of repeat unit A (ortho disposed ether linkages in sub-units) was sulphonated in concentrated sulphuric acid (98% w/w) and the product analysed by 220 MHz nmr spectroscopy as a solution in the concentrated sulphuric acid (the product would not precipitate by pouring the acid solution into water). As with the homopolymer of repeat unit A (para disposed ether linkages in sub-units) the product was found to have sulphonated rapidly with substantially all the sub-units bearing one 35 -SO<sub>2</sub>OH group.

#### EXAMPLE 9

A homopolymer of repeat unit A (meta disposed ether linkages in sub-units) was sulphonated in concentrated sulphuric acid (98% w/w) and the product analysed by 220 5 MHz nmr spectroscopy as a solution in the concentrated sulphuric acid (the product would not precipitate by pouring the acid solution into water). As with the homopolymers of repeat unit A (para or ortho disposed ether linkages in the sub-units) the product was found to have sulphonated rapidly; substantially all the sub-units bore two -SO<sub>2</sub>OH groups however.

# EXAMPLE 10

A homopolymer containing the repeat unit B (50 g) was dissolved in concentrated sulphuric acid (98% w/w)

15 (200 ml), left for 72 hours, precipitated in dilute sulphuric acid, washed, macerated and dried. The infra-red and nmr spectra of the product showed it to be indentical with the starting material.

# EXAMPLE 11

20 A homopolymer containing the repeat unit B (20 g)
was added to chlorosulphonic acid (100 ml). Slight
effeverence was apparent and white fumes were evolved.
After standing overnight (the total time of the polymer in
the acid being 20 hours), the solution was poured into
25 300 ml of concentrated sulphuric acid (98% w/w) and this
solution then poured onto ice. A very fine precipitate
resulted, which filtered very slowly and was washed once
with water and then dried. The product appeared to be
almost entirely soluble in water - indicating a high
30 degree of sulphonation.

## EXAMPLE 12

A homopolymer of repeat unit B (20 g) was mixed with oleum (150 ml) and left overnight. The resulting black solution still contained undissolved polymer and so the .

35 mixture was stirred for a further 5 hours, and then poured

into water. No precipitate was obtained, but instead a brownish/orange solution was obtained. The product was discarded, being assumed to be highly sulphonated or entirely degraded.

- In another experiment, the homopolymer of repeat unit B (15 g) was dissolved in concentrated sulphuric acid (98% w/w); oleum (50 ml) was then added and the solution stirred for 30 minutes and then poured onto ice and left overnight. A rubbery white solid was
- 10 precipitated and was washed and dried. This product was found to be highly sulphonated.

#### CLAIMS

1. A polyarylethersulphone copolymer which is a hydrophilic sulphonated copolymer derived by controllably sulphonating a copolymer having the repeat unit of formula

together with the repeat unit of formula

substantially all the units A being sulphonated after sulphonation and substantially all the units B remaining non-sulphonated after sulphonation.

- 2. A polyarylethersulphone copolymer according to claim 1 derived by controllably sulphonating a copolymer which has 1 to 99 mole % of units A and 99 to 1 mole % of units B.
- 3. A polyarylethersulphone copolymer according to claim 2 derived by controllably sulphonating a copolymer which has 5 to 80 mole % of units A and 95 to 20 mole % of units B.
- 4. A polyarylethersulphone copolymer according to 20 claim 1 having a hydrophilicity corresponding to a water absorbtion capacity at ambient temperature of 2 weight % water absorption to complete solubility in water.
  - 5. A polyarylethersulphone copolymer according to claim 4 having a hydrophilicity corresponding to a water absorption capacity at ambient temperature of 2 to 40 weight %.
    - 6. A polyarylethersulphone copolymer according to any one of the preceding claims in which the sulphonyl groups on the copolymer have the formula  $-SO_2OY$  where Y is H, Cl, an alkali metal or NR<sub>4</sub> where R is an alkyl group.

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- 7. A polyarylethersulphone copolymer according to any one of the preceding claims in the form of a membrane.
- 8. A process for the production of a hydrophilic polyarylethersulphone copolymer which comprises controllably sulphonating using concentrated sulphuric acid (98% w/w) a copolymer having the repeat unit of formula

10 together with the repeat unit of formula

$$\mathsf{B} \qquad \mathsf{-o-} \underbrace{\mathsf{O}} \mathsf{-so}_2 \mathsf{-} \underbrace{\mathsf{O}} \mathsf{-so}_2 \mathsf{-so}_2 \mathsf{-} \underbrace{\mathsf{O}} \mathsf{-so}_2 \mathsf$$

to produce a sulphonated copolymer in which substantially all the units A are sulphonated and substantially all the units B are non-sulphonated.

- 9. A process according to claim 8 wherein the copolymer which is sulphonated has 1 to 99 mole % of units A and 99 to 1 mole % of units B.
- 10. A process according to claim 9 wherein the copolymer which is sulphonated has 5 to 80 mole % of units 20 A and 95 to 20 mole % of units B.
- 11. A process according to any one of claims 8 to 10 wherein after sulphonation with the concentrated sulphuric acid (98% w/w), the -SO<sub>2</sub>OH groups formed on the polymer are converted to groups of formula -SO<sub>2</sub>OM where M is an alkali metal or NR<sub>4</sub> where R is an alkyl group.

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# EUROPEAN SEARCH REPORT

EP 79 301 674.2

	DOCUMENTS CONSIDERED TO BE RELEVANT	CLASSIFICATION OF THE APPLICATION (Int. CL3)		
egory	Citation of document with indication, where appropriate, of relevant passages	Relevant to claim		
A.	US - A - 1 054 707 (J.P. QUENTIN)  * claims 1, 4, column 1, line 64 to column 2, line 5 *	1,8,11	C 08 G 75/23 C 08 G 85/00	
A	US - A - 3 709 841 (J.P. QUENTIN)  * claims 1, 3; column 2, lines 3 to 17 and line 27 *	1,8		
A	HOUBEN-WEYLS "Methoden der organischen Chemie" 4th edition, vol. IX 1955, G. THIEME VERLAG, Stuttgert, * pages 460 and 471 *		C 08 F 8/00 C 08 G 65/00	
A	W. FOERST "Ullmanns Encyklopädie der technischen Chemie" 3rd edition, vol. 14, 1963, URBAN AND SCHWARZENBERG,		C 08 G 75/00 C 08 G 85/00 C 08 J 7/00 C 08 L 81/00	
	München, Berlin  * pages 595 to 597 *		CATEGORY OF CITED DOCUMENTS  X: particularly relevant A: technological background O: non-written disclosure P: intermediate document T: theory or principle underly the invention E: conflicting application D: document cited in the application L: citation for other reasons	
X	The present search report has been drawn up for all claims  [Date of completion of the search	Examin	& member of the same pate family, corresponding documenter	
Place	Berlin 03-12-1979		MARX	